Self-administered versus provider-directed sampling in the Anishinaabek Cervical Cancer Screening Study (ACCSS): a qualitative investigation with Canadian First Nations women

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ABSTRACT

Background While (Pap)anicolaoou screening has helped to decrease cervical cancer incidence in Canada, First Nations women continue to have a higher burden and mortality relative to mainstream populations. Many First Nations women may feel uncomfortable with the invasiveness of this test, contributing to this statistic. Implemented from 2009 to 2015 in 10 Northwest Ontario First Nations communities, the Anishinaabek Cervical Cancer Screening Study (ACCSS) uniquely addressed this Indigenous health inequity through a mixed methods approach.

Objective Our goal was to offer an alternative test which the women could do themselves: human papillomavirus (HPV) testing based on self-sampling. We investigated whether First Nations women preferred HPV self-sampling over healthcare provider (HCP)-administered Pap screening.

Methods Participatory action research informed by the ethical space concept has guided all stages of the ACCSS. We conducted qualitative interviews with 16 HCPs and 8 focus group discussions with 69 female community members followed by a cluster-randomised controlled trial (RCT). Here, we draw on the qualitative field data and an end-of-study community update gathering to disseminate and contextualise research findings. Informant data were evaluated using thematic analysis.

Results We discuss factors influencing participants’ strong preference for HPV self-sampling over physician-conducted Pap screening. Key arguments included enhanced accessibility and more personal control, less physical and emotional discomfort and fewer concerns regarding privacy of test results. For future implementation of HPV self-sampling, study participants emphasised the need for more culturally sensitive education addressing community members of all genders, starting at school, clarifying that HPV causes cervical cancer. Further, HPV infection should be de-stigmatised by accentuating that it affects men and women alike.

Conclusion Here we show that self-sampling in conjunction with community engagement and culturally sensitive education and could be a viable option for underscreened Canadian First Nations women. These informant data echo our previous RCT results.

INTRODUCTION

Cervical cancer can be largely prevented through regular screening with Pap(anicolaoou) cytology as the current standard of care in most countries with gynaecological cancer prevention programmes. However, cervical cancer incidence remains high for vulnerable populations. This may be an effect of underscreening4 associated with healthcare access limitations,2 3 related to travel4 5 and inconsistent service.6 7 In addition, women may not participate because of discomfort or embarrassment regarding the invasive nature of the Pap test,6 8 language barriers9 or cultural sensitivity issues.10 11 There is increasing evidence that women see self-sampling for human papillomavirus (HPV) testing as an option for overcoming these inadequacies12 which may lead to significant improvements in screening rates among underscreened women.13-18

Self-sampling is a feasible and cost-effective alternative for enhancing access to screening.
in underserved rural areas and is perceived as being easy to use, given sufficient information is provided to alleviate women’s concerns about correctly inserting the swab and collecting a sample. As the specificity and accuracy of HPV self-testing is comparable to physician-administered HPV tests, self-sampling has been approved as an option for underscreened or never-screened women in the Netherlands since 2001 and is slated for approval in Australia by 2017. Some concern has been raised about the ability of this method to serve ethnic, religious, educational and age groups that historically have been under-represented in conventional screening programmes. In contrast, initial studies conducted with Indigenous peoples of Canada and the USA, that is, Inuit and First Nations women in Canada as well as Hopi women in the USA, suggest that self-sampling is preferable to provider-directed Pap screening.

In Canada, Australia, New Zealand and the USA, Indigenous women have disproportionately high incidence and mortality rates from cervical cancer. In the Canadian province of Ontario, First Nations women are twice as likely to be diagnosed with cervical cancer and die from it. To address this Indigenous health inequity, we conducted the mixed methods Anishinaabek Cervical Cancer Screening Study (ACCSS) which consisted of qualitative and quantitative components. The ACCSS used a participatory action research (PAR) approach informed by Ermine’s ethical space concept. Our aim was to assess whether offering alternative self-sampling could lead to increased screening participation. Here, we present our qualitative results. We draw on testimonies obtained through interviews and focus group discussions as well as a knowledge transfer gathering at the end of study. These findings endorse our quantitative data from the ACCSS’ randomised controlled trial (RCT) where women were twice as likely to get screened if offered self-sampling rather than Pap testing.

**METHODOLOGY**

**Research context**

The principal investigator (IZ), a non-Indigenous, female cancer biologist (PhD) with a background in cultural anthropology and an interest in Indigenous health disparities to address cervical cancer rates in Ontario, Canada, conducted a pilot study (April to December 2009) with one First Nations community in the Thunder Bay region. The majority of participants (87%) preferred HPV self-sampling over Pap testing. IZ was then invited to the Northern Superior All Chiefs Meeting in October 2010 to expand the screening study to other regional communities. Research agreements were formally ratified (winter 2010 and spring 2011) followed by ‘Meet and Greet’ information sessions conducted in the 10 participating communities (spring 2011) to discuss the research. These Robinson Superior Treaty communities which are part of the Anishinaabek Nation lie within a 500 km radius of the city of Thunder Bay, the region’s health and social service centre. They have on reserve populations ranging from 70 to 832 members.

**Research approach**

PAR informed by Ermine’s ethical space concept (2007) to bridge diverging cultural worldviews has guided the ACCSS design, conduct, data analysis and results’ dissemination with continuous communication between partner communities and the multidisciplinary academic team (medical sciences, eg, cancer biology, epidemiology and virology as well as social sciences, eg, sociology, medical anthropology and women’s studies) (www.accssfn.com). In addition to field trips to the communities for data collection, regular teleconferences have enabled active and ongoing participation of members from more geographically distant communities and helped to link communities and project staff to one another. A publication steering committee (PSC) consisting of one representative from each participating community—that is, the senior health manager, was established to review prospective reports and publications with up to three ad hoc PSC members or delegates recruited as coauthors for each submitted peer-reviewed publication. Each publication was also sent to the communities’ political leadership for approval.

**Study design**

To add rigour to our investigation and to understand the dynamics of cervical cancer screening attitudes, a mixed methods study design was used for the larger study conducted between August 2011 and October 2015. In the qualitative component, with parts reported here, we obtained information in two ways: (1) through in-depth interviews with 16 indigenous community healthcare providers (HCPs) with a broad range of roles (from all 10 communities; summer 2011) and (2) through 8 focus groups with a total of 69 community women (from 8 communities; spring and summer 2012). Thereafter, sample saturation was reached. In the quantitative component, an RCT was conducted with the same 10 partner communities with 834 eligible women to assess uptake of self-administered HPV sampling versus Pap screening (spring 2013 to spring 2014). Together with community-based research assistants (CBRAs)—recruited from women living in the partner communities—IZ conducted educational sessions to explain the study during the RCT. CBRAs enrolled participants and provided self-sampling kits or organised Pap-screening. Participants could choose the HCP who would report results back to them. All women with positive test results were immediately referred to the provincial standard of care for treatment and/or follow-up. Data management and analysis occurred between summer 2014 to summer 2015.
Sampling strategy
Interviews and focus groups
The senior health manager in each community recruited 1–2 HCP informants to participate in individual interviews through purposive sampling. The HCPs approached were those most familiar with community reproductive care and sexual health service provision.40 Their ages ranged from mid-20s to late 60s. Occupations included physicians, nurses, nurse-practitioners, social workers and community health representatives. The length of time they had spent working in their community varied between 6 months and 15 years. Fifteen HCPs were female and one was male. Of these, 10 women self-identified as First Nations. Their intersectional perspectives45 as HCPs, as First Nations peoples and as community residents enriched the interviews and allowed a better understanding of the complex issues impacting the cervical cancer screening environment. Ten interviews were conducted in-person and six via videoconference.

At the recommendation of our community partners, we did focus groups (resembling talking circles) rather than individual interviews with community women to enhance their comfort participating in research on a sensitive topic. The senior community health managers recruited participants of mixed ages, including teens, middle-aged women and elders (up to ~70 years of age). Women who self-identified as First Nations female residents in any of the participating communities with a minimum age of 18 and without formal health education beyond secondary school (for a diversity of perspectives and to complement the voices of HCPs) were eligible. A meal was shared during the sessions (mostly held at community centres) and, as is common for research with indigenous communities in Canada, participants were offered a monetary incentive ($C75), to show respect for their time and contributions to the project. IZ conducted the interviews and focus groups with a medical anthropologist (Dr Marion Maar, Acknowledgements) skilled in leading individual and group interviews. HCP interviews and focus groups followed an open-ended, semistructured grounded theory approach, enabling new ideas and questions to arise from the participants, enriching the researchers’ thinking and concepts. For comparative purposes, key topics were probed in each focus group. Supplementary questions and discussion were guided by participant response and questions to the facilitators. Sessions usually began with a broad stimulus question on the experiences with cancer in their community, followed by community concerns about cancer, knowledge and awareness of cervical cancer screening, availability and experiences with PAP screening, barriers and facilitators to screening, a description of HPV self-sampling and interest in a self-sampling trial (online supplementary material). Interviews lasted 40–90 min and focus groups 80–120 min, with participants’ permission to be audiotaped for later analysis. A note taker was present during focus group sessions. No individual or focus group participant in the 10 communities, reported here, withdrew.

Community update gathering
Our qualitative data set also included reflections on self-sampling versus Pap screening as reported and discussed at the 2-day ACCSS community update gathering (CUG) held in a Thunder Bay hotel at the end of study in October 2015. Altogether 35 persons (32 females and 3 males) were in attendance: partner community representatives (13 HCPs by invitation of 1–2 representatives per community decided by each community’s senior health manager—some of whom had participated in the interviews; 4 CBRAs who helped with recruitment during the RCT42—none had participated in the interviews or focus groups; 1 elder; n=18), 6 ACCSS researchers, 1 commissioned Indigenous organiser and facilitator, 1 Indigenous academic researcher outside the ACCSS team, 5 delegates from Cancer Care Ontario and 1 from the Society of Gynaecologists and Obstetricians of Canada, 1 Anishinaabek male artist, who had designed the ACCSS study logos (www.accssfn.com), 2 local male undergraduate students volunteering for the ACCSS and 3 female note takers. The CUG used multiple discussion formats modelled on talking circles to gather participant perspectives on the successes and challenges of the ACCSS. We began with an opening talking circle which allowed participants to offer their spontaneous reflections on the ACCSS, followed by a World Café structured group discussion format46 in which participants were assigned to one of three groups and moved through three facilitated discussion stations each addressing an assigned question: (1) what worked and what could have been done better, (2) best approaches to deliver culturally sensitive cancer prevention education and (3) future directions based on project successes. After the World Café session, a summative talking circle ended the gathering.

Data analysis
Multiple levels of analysis were employed with the qualitative data. Deidentified transcripts from interviews, focus groups and the CUG were manually coded by IZ and PW using open coding47 to identify basic themes which were grouped into global themes through review and discussion.48 A commissioned rural health senior researcher (Dr Mary Ellen Hill) uploaded the deidentified transcripts into NVivo 9 (QSR International) and assisted us to more comprehensively identify the multiple locations and contexts in which the basic and global themes appeared in the transcripts. We shared our findings and obtained feedback from the community informants via an educational workshop held in Thunder Bay in October 2012 (interviewees),49 community visits from December 2013 to March 2014 (focus group members) and community visits in December 2015 (CUG participants and community leadership). No diverging evidence or inconsistent data were identified. Independent review of the transcripts, code book and discussions by IZ and PW44 led to the identification of 15 basic and 5 global themes (figure 1).
Figure 1  Basic and global themes deduced from interviews and focus groups. STI, sexually transmitted infection.

In the quotes cited below, I stands for interview, FG for focus group, and regarding the CUG, OC for opening (talking) circle, WC1—3 for respective World Café (discussion station) or SC for summative (talking) circle. Only quotes from community representatives were reported. Due to confidentiality reasons, these were not individualised.

RESULTS
Cervical screening practices in partner communities

No regular screening services
Pap testing is the standard of care for cervical cancer screening in Ontario. Screening is opportunistic without a formal recall system and participation rates have not been assessed. As community visits from locum physicians or nurse practitioners are usually infrequent and community nurses are not certified to do Pap testing, many women living in small rural First Nations communities need to travel to nearby towns to attend Pap clinics in conjunction with seeking other services. The time required to travel and keep appointments is a disruption to their everyday lives and a significant barrier to accessing care. As a woman emphasised: ‘Most of us do not have doctors here … you have to drive up to [nearby town]’ (FG1).

Lack of provider flexibility
Those who can access care through nearby small-town health centres and family health teams, located 40–60 km

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away, reported that clinics often only offered Pap tests on certain days of the week. Frequently, physicians were called away to emergencies, so appointments were cancelled and rescheduled. Recounting her experience, a woman said: ‘I don’t know how many times they’ve cancelled Pap tests and [when I asked] “okay well can I get it on Friday” … [heard back] “No, we don’t do them on Friday”’ (FG2).

Transportation and child care issues
Women who elected to access care through a family practitioner, walk-in clinic or gynaecologist in Thunder Bay, located 60–200 km away from their communities, often found that it was difficult to arrange transportation. Space was limited in the community medical van, which travelled to the city only certain days of the week and, if overnight stay was required, accommodations had to be found: ‘If the van was full, [you would] have to get a ride … some of us don’t have cars, you know’ (FG7). Women with children and work commitments also had difficulty arranging childcare and time off work to keep their appointments. As a HCP commented: ‘It certainly does pose challenges with regards to babysitting care and the mother being away for an entire day’ (I16).

No efficient follow-up
Some women noted that they had to ‘wait and wait’ to hear back from their providers and, in many instances, did not hear back about their results until they visited the health centre for another reason. As one focus group participant noted: ‘It’s crazy here, how long you have to wait for anything and half the time, you don’t even get a call … you go to your next doctor’s appointment for something else and [have to ask] … by the way, how about my Pap two months ago?’ (FG2).

Privacy concerns
Those who were recalled for results spoke of privacy concerns at small community health centres where staff ‘phoning people, telling them, ‘we’ve got your results … you need to make another appointment’ (FG2) was upsetting. As one woman said: ‘I don’t want nobody else to know’ (FG2). Due to stigma, they need reassurance that results ‘did not get out into the community’ (I16).

Conversely, there was a view that self-sampling was ‘a lot more private, at home, if you do it by yourself’ (FG4). Picking up a self-test, doing the test at home, mailing it away and getting results by mail offered much more privacy and confidentiality than the alternative of having to make appointments at the health centre. One woman said: ‘Women would prefer to be discreet, do it themselves and get their own results and not have their results shared with others’ (FG1).

Physical comfort level
Paps hurt
Although some women who were used to getting Paps felt that ‘it’s kind of a routine after a while … they open you up and they swab and then you’re done’ (FG2) or ‘the Pap is uncomfortable but it has to be done’ (FG4), other women continued to experience anxiety, even after having Paps on a regular basis: ‘it doesn’t get any easier, like the first time and then the next year, it didn’t get any easier for me’ (FG2). One woman who had gone for one and never had another Pap said: ‘I didn’t like the way it felt, that’s why I didn’t want to go back there’ (FG9). Another said that the ‘pain’ associated with the test had stopped a lot of women: ‘They don’t want their Paps [be] cause it hurt’ (FG1). The test also could be exceptionally painful if providers were ‘in a rush’ or ‘rough’ and did the procedure ‘real quick’ (FG8). From the point of view of an HCP: ‘Some women, when they come to the appointment, they decide they don’t want to get a Pap, because it’s uncomfortable, they’re just afraid’ (I8).

Self-sampling is less painful
Self-sampling was seen as advantageous and may increase screening participation because the procedure itself was thought of as being ‘less invasive’ and ‘less painful’ than having a Pap test (I6). The prospect of testing using a ‘swab’ yourself was much more comfortable than having to submit to an HCP examination with ‘metal or plastic instruments’ (FG1). As a woman said: ‘Women wouldn’t be so agitated and nervous about having the [self-] test’ (FG4).

HCPs also supported the idea of self-sampling, seeing it as less invasive, as this quote suggests: ‘It’s a lot less clinical … stripping down and allowing someone else to do the scraping of the cervix, the whole uncomfortable procedure of going through a Pap opposed to doing it privately in the bathroom on your own is a huge difference. …I think the prevalence [participation] rate would go up’ (I1). The Pap procedure on the other hand was viewed as being ‘such an invasive part of a check-up from the doctor [and] women don’t like to get that done’ (FG9).

Similar opinions were also expressed at the CUG:

[Self-sampling] is less invasive. A lot of them experience … not good memories growing up and what not; so it was a little harder to get the Elders involved, but once we did, it kind of made the women more comfortable and we got a lot of results through the community there. [I said] if you wanted to come by my house … I have the kits, I got a washroom and whatever … It was always on their terms because I felt like giving it to them on their terms was better … It was really, really good for our community because it’s like, uh, the snowball effect … And now I have people asking me like when can I do this [self-sampling] … but a lot of the women shied away from the Pap test. (SC)
practitioner] because maybe she’s going through the same thing that you’re going through’ (FG2). Others felt that they would be ‘embarrassed anyway, no matter who did it’ (FG2). Likewise, a young woman said: ‘I only got a Pap test once and I got it from a woman … and she was trying to make me feel comfortable, but it’s still really awkward’ (FG4).

**Age factor**

There was a view, as an HCP stated that: ‘The older women, and then the really young newly sexually active young girls are a little reluctant to get their Paps. We have a long time with that’ (I1). Young women often delayed making appointments for Pap tests and birth control for fear that their HCPs, and by extension, their families, would know that they were sexually active. This relates to sexual stigma as one woman said: ‘I was sexually active for a while before I even went for my first one … I was nervous about it because … all the nurses and the doctors know each other’ (FG2). Older women with histories of sexual abuse or residential school experiences were reluctant to go to HCPs generally as they were fearful of being ‘touched’ (I17). In the words of a focus group participant: ‘If you’re a culture that has been impacted by residential school, colonization … it’s going to be … ten times harder for us to go to a male doctor’ (FG1). An HCP said: ‘The reason why they don’t go get screened [is] maybe because their suffering from post-traumatic stress and [going for a Pap test] maybe it triggers [the stress] in people’ (I4).

At the CUG, respondents also commented on the interest in self-sampling among young women in their communities, including those under 25 who were not yet eligible to participate. They noted that this was a population to whom education and screening should be extended to capitalise on continuing screening for the future:

> What I want to add, is to have that sustainability, I think we need to address the younger ones to get them ready for 26 and over, you know. Like some communities I work with, I can see that it’s staying around because they still ask me, can we do the self-sample? Sure, they’re all about the self-sampling too because I mean it’s actually based on stuff that [they are comfortable with]. (WC3)

**High comfort with self-sampling**

The consensus was that self-sampling for HPV testing would increase comfort and reduce embarrassment. Women commented that ‘doing it yourself would be better’ than having an HCP (FG2). Others emphasised they ‘would feel more comfortable with that [self-test] than a male doctor’ (FG5). Doing one’s own test was also seen as easier for people who reported being frightened or ‘shy’ of doctors or nurses who visited the communities (FG8). Self-testing would also address the ‘trust issues’ that discouraged women from seeking care from non-indigenous HCPs (FG5).

**Convenience factor self-sampling**

**More privacy**

Having the test available at home might encourage more women to do the test: ‘I think more people would monitor it that way and test it themselves, like ‘well maybe not today, but … eventually I’m going to try it’ (FG2). From an HCP perspective, giving women the option of self-testing in a clinical setting or giving them the option of taking it home might make women ‘feel more empowered’ and contribute to a better relationship with HCPs (I8), or first-time users could get help from their provider: ‘An option, they can either do it themselves, or you can offer to do it for them’ (I4).

**No appointment needed for self-sampling**

Summing up her views regarding self-sampling, a focus group participant said: ‘It is more comfortable to do it at home … it’s simple’. The test would also eliminate the necessity of scheduling out-of-town appointments and travelling to a small town or city to access care (FG6). Another said: ‘I think the self-test is beneficial for them all because sometimes people don’t have time for appointments to take off work, it [the Pap] is kind of an inconvenience’. Self-testing was also viewed as ‘more convenient’ because women could keep the kits at home (FG4).

Overall, the ease of accessing self-sampling seemed appealing to younger women who were busy with work and childcare and ‘would go for it’; less so for some older woman who had fewer constraints on their time and ‘were used to getting Paps’ (FG4). Regarding the convenience of self-testing, an HCP said: ‘Even hearing [about self-testing], people are just, What? Oh, I’d do that, for sure, instead of me going to the doctor’ (I3). Another provider said: ‘Doing a self-test doesn’t take very long … it’s something that can be dealt with, done, gone’ (I10).

The above voices were echoed during the CUG reinforcing that self-sampling is comfortable and enhances their control of the screening process as the following quote illustrates:

> I really strongly believe that … because I was the one that was doing it [self-sampling], I was the one that was in control … and this way it gave me the ability to do it myself and I got all the results, they were fine; … it was also self-empowering, great, I like that. (OC)

**Educational implications**

**Need to explain self-sampling**

While the informants generally reacted positively towards ‘a new screening tool’, some of the HCPs indicated reservations for immediate use of self-testing. HCPs felt that education around self-testing might better be done in the context of a well woman healthcare visit, as ‘part of a physical, that might get best [results]’ (I8). They were concerned that, without education, women would not necessarily understand that they would still need to go to their HCP for other reproductive healthcare: ‘They
would, they’d probably think, well, I’ve had that done, so I don’t need this done’ (I8).

Further emphasising the need for education, one HCP interviewee said: ‘If you hand me a kit, I won’t touch it. Because I wouldn’t know what to do, what if I did it wrong, or whatever, right, so? And I think a lot of people would be that way’ (I3). Another expressed a similar concern: ‘If you just give a test kit to them at home, the majority of them are just going to throw them out’.

**Women being blamed for sexually transmitted infections**

Reluctance to self-test for HPV was also related to fear that women would be blamed for having a sexually transmitted infection (STI) : ‘HPV, like, whoa, I don’t have that, like, I don’t even want to know if I have that’ (I8). Women were distressed about testing for STIs, ‘because it’s a relationship thing’ that implied that someone had ‘cheated’ (I1). Indeed, a focus group participant wondered: ‘… do they [the men] not need to know some of this stuff too?’ (FG1). ‘… I know only women get cervical cancer but … they’re helping it along in the same sense where they’ve passing it back and forth’ (FG1).

**Contribution to stereotype**

Focus group participants were concerned that any publication of information about HPV rates on reserve could have negative effects, contributing to the stereotype that ‘all native people have HPV’ (FG6). As one HCP summarised: ‘A lot more education has to be done about HPV and cervical cancer together’ to alleviate these concerns (I8).

**DISCUSSION**

The ACCSS provides insights into an important Indigenous health inequity as it uniquely addresses high cervical cancer rates in Canadian First Nations using a mixed methods qualitative and quantitative approach. Face-to-face testimonies by our informants, especially those from the focus groups, demonstrate that self-sampling is the preferred method over Pap testing in the ACCSS partner communities—supporting our previous quantitative RCT data.

At the 2-day CUG which was held to discuss the successes and challenges of the ACCSS, the community representatives confirmed a strong preference among the women in their communities for HPV self-sampling. They concluded that offering self-sampling on a continuing basis would maximise women’s ability to screen at times and in places which would make them most comfortable and enhance their control of the screening process. A limitation of our study is that we included only 10 First Nations communities from Northwest Ontario in Canada. However, other Indigenous women worldwide share similar colonial experiences and socioeconomic disadvantages. While our findings require validation, they may be helpful for initiating comparative studies in Canada and elsewhere.

Worldwide, women not participating in Pap testing tend to accept HPV self-sampling instead—a fact largely resonating with the voices from our informants: the self-test has been reported to overcome issues with transportation, childcare and the inconvenience of having to go to a clinic for screening; it increases women’s autonomy and addresses social barriers in rural communities; it is viewed as more private and convenient; and it also alleviates some of the personal stress, for example, physical and emotional discomfort of having Paps. Those who prefer Paps have more confidence in physicians because they are concerned about doing the self-test accurately, This and the concern that women may not show up for other reproductive healthcare when doing self-sampling were raised by a few HCP informants which, as also suggested by them, could be addressed through appropriate education prior to testing.

To make women aware of the benefits of cervical screening through culturally sensitive education was a recurring theme throughout the process of the ACCSS and has not been testified as such in other studies. Our informants recommended educational strategies adapted to their communities’ needs, various age groups and both sexes as previously reported by us. To make HPV self-sampling successful, appropriate information, such as is provided along with the faecal blood self-test for colon cancer, is necessary, as also pointed out in other research. To de-stigmatise sexually transmitted HPV, men must be made aware that both they and women can be carriers of HPV infecting each other. The current notion, suggested by our informants, is that men in their communities are unaware of this fact. This may contribute negatively to younger women attending cervical screening. This limitation in information still needs to be adequately addressed as cervical screening generally is scheduled around sexual activity (often without HCPs mentioning the cause) and self-sampling is valid for HPV testing only.

In conclusion, most ACCSS informants communicated a clear preference for HPV self-sampling over Pap screening. Offering this alternate cervical screening tool would free up HCP time and eliminate several structural barriers. HPV testing is a superior proxy over Pap cytology to detect cervical lesions early enough for treatment and has already become the primary cervical cancer check in some countries. The ACCSS results should be helpful to stakeholders and decision makers who are in charge to design future screening programme including follow-up of abnormal results and access to quality care. Special requirements for underserved populations are required to overcome current barriers discussed here and previously.

The ACCSS team regularly shares results with its Anishinaabek partner communities, the public, decision makers and the scientific community. The end of study CUG was an example of such knowledge dissemination. Overall, and most likely due to the PAR approach, our experience with the Anishinaabek partner communities has been truly positive with a fruitful and candid
exchange of ideas between both groups. This became increasingly clear during the CUG with a reflection on lessons learnt and strong community-guided suggestions for future education and research (Wakewich and Zehbe, in preparation).

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Competing interests None declared.

Patient consent Obtained.

Ethics approval The ACCSS was approved by the principal investigator’s host institution (Lakehead University Research Ethics Board #038 11-12/ROMEO #1463139 and 12-13/ROMEO #1463139). As requested for ethics approval, the ACCSS design is based on the Ownership, Control, Access and Possession (OCAP) Guidelines recommended by the First Nations Information Governance Centre (FNIGC, 2016; http://fnigc.ca/ocap.html) and respects the terms of the tri-council policy statement on research involving the first nations, inuit and metis people of Canada (CIHR, NSERC, SSHRC 2014; http://www.pre.ethics.gc.ca/eng/policy-politique/initiatives/tpcsp2-epct2/chapter9-chapit9/).

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